# **circRPS19 affects HK2‑mediated aerobic glycolysis and cell viability via the miR‑125a‑5p/USP7 pathway in gastric cancer**

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**Abstract.** Despite advances in diagnosis and treatment, gastric cancer (GC) remains a refractory disease, which limits overall survival. Therefore, it is key to identify novel targets to develop more effective and precise treatment. Circular RNAs (circRNAs) serve essential roles in the process of various human cancers. Through analyzing GSE83521 dataset, the present study identified a novel circRNA derived from ribosomal protein S19 (circRPS19), which was considered a potential treatment target for GC. Results of RT‑qPCR indicated that circRPS19 was upregulated in GC compared with normal gastric epithelial cells. Loss‑of function assays revealed that silencing of circRPS19 suppressed proliferation and aerobic glycolysis but increased apoptosis of GC cells. circRPS19 upregulated ubiquitin‑specific processing protease 7 (USP7) expression by sponging microRNA (miR)‑125a‑5p. circRPS19 stabilized hexokinase 2 (HK2) protein by USP7‑mediated deubiquitination of HK2. *In vivo* experiments confirmed that circRPS19 promoted GC progression and aerobic glycolysis. Taken together, circRPS19 induced aerobic glycolysis of GC cells by stabilizing HK2 protein via the miR‑125a‑5p/USP7 axis and thus promoting the progression of GC. These findings suggested that circRPS19 served a critical role in the progression of GC and may be a novel therapeutic target for GC.

# **Introduction**

As a malignancy of the digestive system, gastric cancer (GC) originates in the gastric mucosal epithelium (1). Based on global cancer statistics in 2020, more than one million patients were newly diagnosed with GC and ~769,000 patients died from GC; GC ranks fifth in incidence and fourth in

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cancer‑associated mortality of all types of cancer (2). The occurrence and development of GC are influenced by multiple factors, such as diet, family history, consumption of alcohol and tobacco and infection of *Helicobacter pylori* and Epstein‑Barr virus(3). Endoscopic ultrasound is the most efficient diagnostic methods for early‑stage GC, however, a majority of patients are diagnosed at advanced stages (4). Surgical resection is mainly applied to treat early-stage patients, while chemotherapy proves most efficient in advanced‑staged patients (5). In spite of improvement in GC diagnosis and treatment, the prognosis of GC still remains poor and the 5‑year survival rate is <30% (6). Hence, it is of great importance to identify novel molecular mechanisms in GC progression.

Circular RNA (circRNA) is a type of RNA with covalently closed structure, generated by back‑splicing of precursor mRNA (7,8). Former studies have revealed the effect of circRNAs in GC progression: For examples, circRNA\_100290 modulated by eukaryotic translation initiation factor 4A3 (EIF4A3) can interact with microRNA (miRNA or miR)‑29b‑3p to target integrin subunit alpha 11 (ITGA11), thus facilitating proliferation and invasion of GC cells (9); circRNA nuclear receptor interacting protein 1 targets the Akt1/mTOR pathway by sequestering miR‑149‑5p to promote GC progression (10); circRNA ELAV like RNA binding protein 1 (circ-HuR) downregulates HuR expression by binding to CCHC‑type zinc finger nucleic acid binding protein (CNBP), thereby suppressing metastasis, proliferation and invasion of GC cells (11); the circ\_0004872/miR‑224/Smad4/ADAR1 axis forms a negative feedback loop to hinder GC progression (12). A recent study showed that *Helicobacter pylori* infection‑induced circRNA mannosidase alpha class 1A member 2 (circMAN1A2) promotes GC progression via sponging miR‑1236‑3p (13). The aforementioned findings indicate that circRNAs are crucial regulators in GC development, highlighting the research value of circRNAs in GC. The present study investigated the role of a new‑found GC‑associated circRPS19 selected from Gene Expression Omnibus (GEO) database in GC cell proliferation.

Aerobic glycolysis, or the Warburg effect, was discovered by Otto Warburg in the 1920s. Tumor cells, despite sufficient oxygen, prefer the conversion of glucose into lactate to generate energy (14). As reported by previous studies, aerobic glycolysis serves a key role in a variety of carcinomas, including GC: For example, long non‑coding RNA H19 (imprinted maternally expressed transcript interacts with miR‑519d‑3p to regulate

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lactate dehydrogenase A (LDHA), thus facilitating aerobic glycolysis and proliferative ability of GC cells (15); Opa interacting protein 5 antisense RNA 1 (OIP5‑AS1) sequesters miR‑186 to enhance proliferation and aerobic glycolysis in GC cells (16); pyruvate kinase M2 (PKM2) serves an oncogenic role in GC cells by facilitating aerobic glycolysis (17); Hexokinase 2 (HK2), as a rate-limiting enzyme, catalyzes glucose metabolism and cancer cells commonly exhibit increased glucose metabolism (18). Previous research has demonstrated that circRNA functions as a regulator of HK2 in GC cells to facilitate aerobic glycolysis (19). Nevertheless, the association between HK2 and circRPS19 in GC cells needs to be further investigated.

The present current study aimed to investigate the functions of circRPS19 in GC cell proliferation and aerobic glycolysis and its downstream molecular mechanism.

# **Materials and methods**

*Cell culture and treatment.* Human GC cell lines (AGS, HGC-27, NCI-N87 and SNU-1) were obtained from China Center for Type Culture Collection and normal human gastric epithelial cell line (GES‑1) was obtained from American Type Culture Collection (ATCC). All cell lines were cultured in RPMI-1640 medium containing 10% fetal bovine serum (FBS; Gibco; Thermo Fisher Scientific, Inc.). Cell lines were incubated in a humid atmosphere with  $5\%$  CO<sub>2</sub> at  $37^{\circ}$ C.

For RNA stability analysis, GC cells were treated with 3 U/µg RNase R (9001‑99‑4, Sigma‑Aldrich; Merck KGaA) for 24 h at room temperature or treated with 2  $\mu$ g/ml ActD (50‑76‑0, Merck) for different time intervals (0, 6, 12, 18, 24 h) at room temperature. RNA stability was determined by using RT‑qPCR as following description.

To analyze protein stability, GC cells were treated with  $5 \mu$ M cycloheximide (CHX, HY-12320, MedChemExpress) for 0, 3, 6, 9 h or treated with 20  $\mu$ M MG132 (HY-13259, MedChemExpress) for 4 h. Cells were treated with 10  $\mu$ M HBX19818 (HY‑17540, MedChemExpress) to inhibit USP7 expression. Protein levels were detected using western blot as following description.

*Data collection and bioinformatics analysis.* circRNAs significantly overexpressed in GC tissues (under the condition of P≤0.05) were screened out from GSE83521 dataset in Gene Expression Omnibus (GEO; ncbi.nlm.nih.gov/gds). starBase database (rnasysu.com/encori/) was used to screen miRNAs that potentially interact with both circRPS19 and USP7.

*Cell transfection.* Small interfering RNAs (siRNAs) targeting circRPS19 or ubiquitin‑specific processing protease 7 (USP7) sequence were synthesized for knockdown with non-targeted siRNA as the negative control (NC). For overexpression of USP7, the full-length sequence of USP7 was inserted into pcDNA3.1 vector to establish USP7 overexpression vector, with empty vector as NC. miR-125a-5p mimics was synthesized and used to upregulate miR‑125a‑5p expression, with mimics-NC as NC. All plasmids and vectors were synthesized and constructed by Guangzhou RiboBio Co., Ltd. Cell transfection was performed at 37˚C using Lipofectamine 2000 (Invitrogen; Thermo Fisher Scientific, Inc.). After being transfected with siRNAs and pcDNA3.1 vectors for 48 h or transfected with for miRNA mimics or inhibitor for 24 h, cells were harvested for subsequent experiments. The duration between end of transfection of siRNAs and pcDNA3.1 vectors and subsequent experiments was 48 h. The duration between end of transfection of miRNA mimics or inhibitor and subsequent experiments is 24 h. siRNA and pcDNA3.1 expression vector  $(0.8/50 \mu l)$  were treated with Lipofectamine 2000 for transfection at 24 well plates. miRNA mimics were transfected at a final concentration of 100 nM. The sequences were as follows: si-NC, 5'-CCGGCGGCCGCAAATTAC GAATGACTCGAGTCATTCGTAATTTGCGGCCGTTTTT G‑3'; si‑circRPS19‑1: 5'‑CCGGCTCATTCGTAACGGCCG CAAACTCGAGGAGTAAGCATTGCCGGCGTTTTTTTT G‑3'; si‑circRPS19‑2: 5'‑CCGGATTCGTAACGGCCGCAA ACTGCTCGAGTAAGCATTGCCGGCGTTTGACTTTTT G‑3'; si‑circRPS19‑3: 5'‑CCGGGCCTCATTCGTAACGGC CGCACTCGAGCGGAGTAAGCATTGCCGGCGTTTTTT G‑3'; mimics‑NC: 5'‑UCUCCGAGAUCUUACCACCUG GUA‑3'; miR‑125a‑5p mimics: 5'‑UCCCUGAGACCCUUU AACCUGUGA‑3'; inhibitor NC: 5'‑UCAUCUCAGGAA GGGUAUCGAGGA‑3'; miR‑125a‑5p inhibitor: 5'‑UCACAG GUUAAAGGGUCUCAGGGA‑3'; si‑NC: 5'‑CCGGGTTAA TTTATTATCCAGCATTCTCGAGAATGCTGGATAATAA ATTAACTTTTTG‑3'; si‑USP7‑1, 5'‑CACCGTGTAATT TATTATCTTCAGCACTCGAGTGCTGAAGATAATAAAT TACA‑3'; si‑USP7‑2, 5'‑CACCATTCATAGAACACTCTT TGTACTCGAGTACAAAGAGTGTTCTATGAA-3'; si-USP7‑3, 5'‑CACCAGTTGTAGATGTAACACTGGTCT CGAGACCAGTGTTACATCTACAAC‑3'.

*Reverse transcription‑quantitative (RT‑q) PCR.* Total RNA was extracted from AGS, HGC‑27, NCI‑N87, SNU‑1 or GES‑1 cells using TRIzol (Invitrogen; Thermo Fisher Scientific, Inc.). RT was performed using the PrimeScript 1st Strand cDNA Synthesis kit (Takara Bio, Inc.) according to the manufacturer's protocol. RT‑qPCR was performed on ABI 7500 fast PCR System with SYBR green PCR Master Mix (both Applied Biosystems; Thermo Fisher Scientific, Inc.). The qRT-PCR reaction was performed 95°C for 5 min, followed by 40 cycles of 95°C for 10 sec and a primer-specific annealing temperature of 60˚C for 30 sec. GAPDH and U6 were used as internal controls. Relative RNA expression was calculated using  $2^{-\Delta\Delta Cq}$  method, as previously described (20). Additionally, gel electrophoresis was performed to identify the circular structure of circRPS19. Briefly, circular structure of circRPS19 was validated by PCR reactions using the HotStar HiFidelity Polymerase Kit (#202602; Qiagen) by using pairs of both convergent and divergent primers to amplify circRPS19. Convergent primers were all designed within the same exon. Both cDNA and genomic DNA (gDNA) were applied as the template for PCR analysis. The sequences for primers were as follows (5'~3'): divergent primer for circRPS19 (forward): CAGCTGCCAACAAGAAGCAT; divergent primer for circRPS19 (reverse): CTGTCCGGCGATTCTGTCC; convergent primer for circRPS19 (forward): ACTGACACCTCA GGGACAAAG; convergent primer for circRPS19 (reverse): ATGAGGCAATTTATTAACCCAGCA; RPS19 (forward): GAACCAGCAGGAGTTCGTCA and reverse): AGAACC AGTTCTCATCGTAGGG; miR‑125a‑5p forward): GCC

GAGTCCCTGAGACCCTTTA; miR‑125a‑5p (reverse): CTC AACTGGTGTCGTGGA; USP7 (forward): CCGAGGACA TGGAGATGGAA and reverse, TTTCGCACAAAACAC GGAGG; GAPDH (forward): GACAGTCAGCCGCATCTT CT; GAPDH (reverse): GCGCCCAATACGACCAAATC; U6 (forward): TCCCTTCGGGGACATCCG; U6 (reverse): AAT TTTGGACCATTTCTCGATTTGT.

*Western blotting.* Total protein was isolated from AGS cells with RIPA lysis buffer (Sigma‑Aldrich; Merck KGaA), Protein concentration was measured by using a bicinchoninic acid (BCA) Kit (Pierce Biotechnology, Rockford, IL, USA). Protein samples (30 ng/lane) were then separated by 10% SDS-PAGE (Bio‑Rad Laboratories, Inc.). Afterwards, the samples were transferred onto PVDF membranes, which were blocked with 5% defatted milk in 5% TBST at room temperature for 2 h. Subsequently, membranes were incubated with primary antibodies, including anti‑HK2 (1/1,000, ab209847, Abcam, MA, USA), anti-6-phosphofructokinase subunit alpha (anti-PFK1, 1/1,000, ab154804, Abcam), anti‑PKM2 (1/1,000, ab85555, Abcam), anti‑USP7 (1/1,000, ab264422, Abcam) and the internal reference anti‑β‑actin (1:1,000, ab8227, Abcam) at 4˚C overnight. After being washed in TBST buffer, the membranes were incubated with horseradish peroxidase (HRP)-conjugated secondary antibodies (1/2,000, ab154804, Abcam) at room temperature for 1 h. Pierce ECL Plus Substrate (Pierce; Thermo Fisher Scientific, Inc.) was then used to visualize blots. Image J software v1.8.0 (National Institutes of Health, USA) was used to quantify protein blots.

*Cell Counting Kit‑8 (CCK‑8) assay.* Transfected AGS or HGC‑27 cells were inoculated into 96‑well culture plates at a density of  $3x10^3$  cells/well. GC cells were incubated with  $10 \mu$ l CCK‑8 solution (Dojindo Laboratories, Inc.) for 2 h. After incubation, the absorbance at 450 nm was detected at different time points (0, 24, 48 and 72 h) by a spectrophotometer (Thermo Fisher Scientific, Inc.).

*EdU assay.* Transfected AGS or HGC‑27 cells were plated in 12-well plates at a density of  $1x10<sup>5</sup>$  cells/well. GC cells were treated with EdU reagent (Guangzhou RiboBio Co., Ltd.). Following incubation at 37°C for 2 h, cells were washed in PBS and fixed in 4% paraformaldehyde (Sangon Biotech Co., Ltd.) at  $4^{\circ}$ C for 30 min. The nuclei of GC cells were stained by  $400 \mu$ l DAPI (Sigma-Aldrich; Merck KGaA) at room temperature for 30 min. GC cell proliferation was observed under a Leica GmbH fluorescence microscope (x200) and quantified by calculating the percentage of EdU‑positive cells in five randomly selected view fields using Image J software v1.8.0 (National Institutes of Health, Bethesda, MD, USA).

*TUNEL assay.* The apoptosis of transfected AGS or HGC-27 cells was measured by TUNEL assay. GC cells were fixed in 4% paraformaldehyde (Sangon Biotech Co., Ltd.) at 4˚C for 30 min. The fixed cells were then washed with 5 ml cold 70% ethanol, and the dehydrated cells were incubated at ‑20˚C for 4 h. Cells were permeabilized with 20  $\mu$ g/ml Proteinase K in PBS for 5 min. Cells were then treated with the ProLong Antifade Kit (P‑7481, Molecular Probes) and incubated with TUNEL reaction mixture (Roche Diagnostics) containing 200 U/ml TdT and 4  $\mu$ M dUTP at room temperature for 1 h. After removing the reaction buffer, cells were washed with PBS for 3 times. GC cell nuclei were stained by  $400 \mu$ l DAPI (Sigma‑Aldrich; Merck KGaA) that was added to the media (1:10) at room temperature for 30 min. The percentage of TUNEL‑positive cells in five randomly selected view fields was observed under a Leica GmbH fluorescence microscope (x200) and calculated using Image J software v1.8.0 (National Institutes of Health, Bethesda, MD, USA).

*Flow cytometry.* Flow cytometry analysis was used to detect cell apoptosis. Briefly, cells were collected and washed twice in cold PBS. Cells were then fixed with 4% paraformaldehyde (Sangon Biotech Co., Ltd.) at ‑20˚C overnight. GC cells were suspended in 600  $\mu$ l eBioscience™ flow cytometry binding buffer (Invitrogen; Thermo Fisher Scientific, Inc.) at a concentration of 10<sup>6</sup> cells/ml and then stained with 5  $\mu$ l Annexin V/FITC and  $5 \mu l$  propidium iodide (both BD Biosciences) in dark for 15 min. Samples were analyzed using a FACSAria flow cytometry (BD Biosciences, San Jose, CA, USA). Results were analyzed using FlowJo 7.6 software (Tree Star, Inc.).

*Measurement of glucose consumption and lactate production.*  The glucose consumption and lactate production of transfected AGS or HGC‑27 cells were detected by using a Glucose Assay kit (CBA086) and a Lactate Assay Kit (MAK064; both Sigma‑Aldrich) as previously described (22).

*Examination of extracellular acidification rate (ECAR) and oxygen consumption rate (OCR).* ECAR and OCR were examined as previously reported (22). Seahorse XF96 (cat. no. 102601‑100, Seahorse Bioscience) was applied for the analysis of ECAR and OCR in AGS or HGC‑27 cells. For ECAR analysis, GC cells were exposed to  $1 \mu$ M oligomycin for elimination of mitochondria ATP production and 150 mM 2‑Deoxy‑D‑glucose (2‑DG) at 37˚C for 1 h for inhibition of aerobic glycolysis (both Sigma‑Aldrich; Merck KGaA) successively, followed by assessment using Glycolytic Stress Test kit (Seahorse Bioscience). For OCR analysis, GC cells were exposed to 1.0  $\mu$ M oligomycin, 0.5  $\mu$ M carbonyl cyanide 4‑(trifluoromethoxy) phenylhydrazone (FCCP; a protonophore uncoupler) and 0.5  $\mu$ M antimycin A at 37°C for 1 h for inhibition of electron transfer chain (all Sigma‑Aldrich; Merck KGaA) successively, then measured by XF Cell Mito Stress kit (Seahorse Bioscience). ECAR and OCR were measured by normalizing to total protein content.

*Fluorescence in situ hybridization (FISH).* In brief, AGS cells were washed in PBS, followed by fixation in 4% paraformaldehyde (Sangon Biotech Co., Ltd.) at 4˚C for 30 min and permeabilization in 0.5% Triton X‑100 (Beijing Leigen Biotechnology Co., Ltd.). Cells were blocked with 50  $\mu$ l goat serum (Sigma-Aldrich; Merck KGaA) for 1 h at room temperature. Primer sets were designed based on the sequence of circRPS19 and USP7 in the pGEM‑T Easy vector (Promega). Digoxigenin‑labeled DNA probes targeting circRPS19 (genomic position: chr19:42373768‑42375484; accession no. NM\_001022) and USP7 mRNA (genomic position: chr16:8985951‑9057341 accession no. NM\_003470.3) or control probes synthesized by Guangzhou RiboBio

Co., Ltd. DNA probes were denatured at 75˚C for 5 min. Afterwards, the probes were incubated with GC cells in 10 µl hybridization buffer (30% formamide, 0.9 M NaCl, 20 mM Tris-HCl at pH 8.0, 0.01% SDS, a  $1-\mu$ M probe, and MQ water) at 55˚C for 4 h. To remove excess probes, samples was washed in 50 ml tubes of washing buffer without salt [20 mM Tris‑HCl, 5 mM EDTA and 0.01% (w/v) SDS] or with 1.8 M NaCl at 47°C for 3 min.

DAPI (Sigma-Aldrich; Merck KGaA) was applied to counterstain nuclei of GC cells at room temperature for 30 min. A confocal microscope (x200, Zeiss AG) was employed to obtain images. The probe sequence for circRPS19: TTTGCGGCC GTTACGAATGA.

To identify the co‑location of USP7 and HK2 proteins, we used the anti-HK2 (1/1,000, ab209847, Abcam, MA, USA), and anti‑USP7 (1/1,000, ab264422, Abcam) to incubated cell lysates at 4°C overnight, followed by incubation with horseradish peroxidase (HRP)‑conjugated secondary antibodies (1/2,000, ab154804, Abcam) at room temperature for 1 h. Results were analyzed using Image J software v1.8.0 (National Institutes of Health).

*Co‑immunoprecipitation (Co‑IP) assay and mass spectrometry.* Co‑IP assay followed by mass spectrometry was used to identify the interacting protein of HK2. Cells were washed twice with pre-cold PBS. Cell lysates were obtained with RIPA lysis buffer (Sigma‑Aldrich; Merck KGaA). For each IP reaction, 50  $\mu$ l lysates were used. After being centrifuged at 14,000 g for 15 min at  $4^{\circ}$ C, superna– tants were collected. The supernatants were incubated with 50  $\mu$ g of the negative control anti-Flag (1/200, 66008-3-Ig; Proteintech) or 50  $\mu$ g Flag-labeled anti-HK2 (1/200, ab209847, Abcam) at 4˚C overnight, then with protein A/G magnetic beads (Thermo Fisher Scientific) in accordance with manufacturer's protocol. After being centrifuged at  $3,000 \text{ x g}$  for 3 min at 4°C, the magnetic beads were deposited at the bottom of the tube. Removing the supernatant, the magnetic beads were washed three times in 1 ml lysis buffer. After being added 15  $\mu$ l 2xSDS loading buffer, the magnetic beads were boiled for 5 min. The gel bands were excised for in‑gel trypsin digestion. Subsequently, extracted peptides were analyzed using nano‑LC‑MS/MS (AB SCIEX TripleT OF 5600) for mass spectrometry. Mass spectrometry weas conducted on an Orbitrap Fusion Tribrid mass spectrometer (Thermo Fisher Scientific, Inc.) equipped with an Easynano UHPLC, using a 75  $\mu$ m x 250 mm Pepmap RSLC reverse phase column with a PepMap 1000 trapping column (Thermo Fisher Scientific). The mass spectrometer was equipped with an electrospray ionization (ESI) and operated in the positive ion mode to monitor the m/z transitions for all peptides and IS. Two multiple reaction monitorings were simultaneously monitored for each peptide using manually optimized declustering potentials (DP), collision energies (CE), collision cell entrance (CEP) and exit potentials (CXP). Nitrogen gas was desorbed from the SPME fibre in the injector port at 200˚C, under splitless injection mode. The spray voltage was set as 5,000 V, the atomization pressure was set as 45 psi, the auxiliary gas pressure was set as 10 psi. Nitrogen gas with high purity (≥99%) was used as the carrier gas (constant flow of 1.1 ml/min; average velocity 37 cm/s).

*IP.* To detect ubiquitination of HK2, transfected AGS cells were cultured for 24 h at room temperature in PBS with 0.1% formaldehyde. Cells were treated with the proteasome inhibitor MG132 (10  $\mu$ M) at room temperature for 8 h. GC cells were lysed in IP buffer (Beyotime Institute of Biotechnology, Shanghai, China) with protease and protein phosphatase inhibitors (Roche Applied Science). Cell lysates (100 ml) were then immunoprecipitated with anti-β-actin (1/1,000, ab8227, Abcam) and anti‑HK2 (1/1,000, ab209847, Abcam) at 4˚C overnight. To measure HK2 ubiquitination, the samples were incubated with anti-ubiquitin (1/1,000, 10201-2-AP, Proteintech Group, Inc.) 4˚C overnight and then subjected to western blot analysis as described above.

*Luciferase reporter assay.* The wild‑type or mutant (MUT) sequences of circRPS19/USP7 3' untranslated region (UTR) were inserted into pmirGLO vectors (Promega Corporation) to construct pmirGLO‑circRPS19/USP7‑3' UTR or pmirGLO‑circRPS19/USP7‑3' UTR (MUT). pmirGLO-circRPS19 and pmirGLO-circRPS19 (MUT) were co-transfected into AGS or HGC-27 cells with miR-125a-5p mimics or mimics‑NC using Lipofectamine 2000 Transcription Reagent (Invitrogen; Thermo Fisher Scientific, Inc.) at 37˚C for 36 h. pmirGLO‑USP7‑3' UTR or pmirGLO‑USP7‑3' UTR (MUT) were co-transfected into AGS or HGC-27 cells with miR‑125a‑5p mimics or mimics‑NC using Lipofectamine 2000 Transcription Reagent (Invitrogen; Thermo Fisher Scientific, Inc.) at 37˚C for 48 h. The duration between end of transfectioin and activity measurement is 48 h. After that, relative luciferase activity was measured using a Dual-Lucy Assay kit (Progema) with *Renilla* luciferase activity as the internal reference. The sequence of miR-125a-5p mimics: 5'‑UCCCUGAGACCCUUUAACCUGUGA‑3'; the sequence of NC‑mimics: 5'‑UCUCCGAGAUCUUACCACCUGGUA‑3'.

*RNA pulldown assay.* RNA pulldown assay was performed in AGS or HGC-27 cells to demonstrate RNA interaction. Briefly, 1  $\mu$ g biotin-labeled circRPS19/USP7-3' UTR synthesized by Guangzhou RiboBio Co., Ltd. was added to 1% Triton X‑100, followed by heating at 95˚C for 2 min and cooling in an ice bath for 3 min. Next, the denatured biotinylated RNA was mixed with 15  $\mu$ l streptavidin beads (Thermo Fisher Scientific, Inc.). RNA‑bead complexes were incubated with 100 ml cell lysate obtained using RIPA lysis buffer. After purification, RNA in pulldown products was analyzed by RT‑qPCR as described above.

*RNA‑binding protein IP (RIP) assay.* AGS cells were lysed in RIPA buffer for RIP assay using Magna RIP kit (MilliporeSigma). Briefly, cell lysate was treated with RIP buffer at room temperature for 3 h and added to 10 mg/ml magnetic beads (Thermo Fisher Scientific, Inc.) conjugated with anti-argonaute 2 (Ago2, 1/200, 04–642, MilliporeSigma) or NC anti-IgG (1/200, AP113, MilliporeSigma). After washing the beads with 500  $\mu$ l RIP Wash Buffer, the complexes were incubated with 150  $\mu$ l proteinase K at 55°C for 30 min to remove protein. The enrichment of circRPS19 or USP7 was analyzed by RT‑qPCR as described above.

*Animal experiments.* A total of 20 male BALB/C nude mice aged 4-6 weeks and weighed  $20±2$  g were obtained from

Shanghai Lab Animal Research Center. All mice were raised in specific pathogen free (SPF) condition at 22‑24˚C with a stable humidity  $(55\pm15\%)$  with free access to food/water in a 12/12 h light/dark cycle, according to the guidelines of the local Animal Care and Use Committee at Affiliated Hospital of Nanjing University of Chinese Medicine. Animal experiments were approved by Ethics Committee of Affiliated Hospital of Nanjing University of Chinese Medicine (approval NO. 2021DW‑35‑01, Nanjing, China). All nude mice were randomly classified into two groups (n=10). Each mouse was subcutaneously injected with  $2x10<sup>6</sup>$  AGS cells transfected with si-NC or si-circRPS19-1 (5 nmol/injection). Tumors were monitored on day 7 after injection and measured every three days. On day 28, all nude mice were subjected to euthanasia via cervical dislocation following intravenous injection with 30 mg/kg pentobarbital sodium (23) and tumors were resected and weighed. Death was confirmed by lack of spontaneous breathing for 2‑3 min and loss of eye blink response. Body weight decrease of 15-20% was taken as the humane endpoint. No animals were prematurely sacrificed according to this endpoint. Tumors were cut into  $4-\mu m$  thick small pieces. The tissues were fixed with 10% formalin for 24 h at room temperature, and then embedded in paraffin at 58‑62˚C for 30 min. The sections were deparaffinized in xylene and rehy‑ drated in a graded series of alcohol. A steamer was used for 20 min in an EDTA buffer for antigen retrieval at 92˚C. Next, samples were treated with 3% hydrogen peroxide in methanol at room temparature for 20 min to block endogenous peroxidase activity and the reaction was blocked by incubating in a 5% solution of non-fat milk for 1 h. Slides were then incubated with polyclonal rabbit antibodies against Ki67 (1:200; cat. no. ab16667, Abcam), HK2 (1/500; cat. no. ab209847, Abcam), or USP7 (1/200, ab239936, Abcam) at room temperature for 1 h, followed by incubation with the secondary antibody HRP-polymer anti-rabbit (1/1,000, ab6721, Abcam) at 37°C for 15 min. The samples were counterstained with DAB) at room temperature for 5 min. Images of five typical visual fields of each section were obtained under a light microscope (x200, DMI1, Leica, Wetzlar, Germany) and analyzed using software (Image‑Pro Plus 6.0; Media Cybernetics, Inc.).

*Statistical analysis.* Unpaired Student's t test was used to analyze differences between two groups. Data from three independent assays are presented as the mean  $\pm$  SD. SPSS v22.0 software (IBM Corp.) was used for data analysis. One‑way ANOVA followed by LSD (one control group) or Tukey (more than one control group) post hoc test was used for comparing >2 groups. P<0.05 was considered to indicate a statistically significant difference.

# **Results**

*CircRPS19 has circular characteristics and is upregulated in GC cells.* circRNAs significantly overexpressed in GC tissue were screened from GEO database. According to the GSE83521 dataset, under the condition of P≤0.05, the top eight circRNAs with the highest logFC values were screened, including circRNA\_102614, circRNA\_103122, circRNA\_102191, circRNA\_100984, circRNA\_102615, circRNA\_102777, circRNA\_102561 and circRNA\_103559 (Fig. 1A). Of these, hsa\_circRNA\_103122 (24), hsa\_circRNA\_102191 (25) and hsa\_circRNA\_102615 (26) have been reported in GC and were excluded from subsequent study. RT‑qPCR was used to probe the expression levels of the other five candidates in human GC (AGS, HGC-27, NCI-N87 and SNU-1) and normal gastric epithelial cell lines (GES‑1). circRNA\_102561 was most significantly upregulated in all four GC cell lines compared with GES-1 (Fig. 1B) and was therefore selected for subsequent experiments. circRNA\_102561 (circRPS19) originates from exons 5 to 6 of its host gene RPS19 (Fig. 1C). PCR-agarose gel electrophoresis and RT‑qPCR were performed in GES‑1 cells to validate the circular structure of circRPS19. As shown by PCR‑agarose gel, circRPS19 was amplified by convergent and divergent primer in cDNA but only amplified by convergent primer in gDNA (Fig. 1D). Moreover, after the treatment of RNase R, circRPS19 expression was almost unchanged, while linear RPS19 mRNA was nearly completely digested. Expression levels of circRPS19 and RPS19 mRNA at 0, 6, 12, 18, 24 h in GES‑1 cells treated with Actinomycin D (ActD) were assessed. circRPS19 expression remained steady, while RPS19 mRNA level decreased with time (Fig. 1F). Collectively, the aforementioned results showed that circRPS19 had a circular structure and was associated with GC.

*circRPS19 knockdown inhibits GC cell proliferation and aerobic glycolysis*. As aforementioned, circRPS19 was upregulated in GC cell lines but its specific effects on the malignant phenotype of GC cells is unknown. It was investigated whether knockdown of circRPS19 alters biological processes of GC cells. RT‑qPCR was used to determine the efficiency of circRPS19‑1‑specific siRNAs on silencing circRPS19 in AGS and HGC-27 cells (Fig. S1A). si-circRPS19-1 and si-circRPS19-2 were selected for subsequent assays owing to their higher efficiency. CCK‑8 and EdU assay showed that GC cell proliferation decreased following circRPS19 knockdown (Fig. 2A and B). TUNEL assay and flow cytometry analysis of GC cells showed that GC cell apoptosis was enhanced following circRPS19 depletion (Fig. 2C and D). circRPS19 facilitated GC cell proliferation but prevented GC cell apoptosis (Fig. 2A‑D). Furthermore, the effect of circRPS19 knockdown on glucose consumption and lactate production of GC cells was assessed. Glucose consumption and lactate production were decreased following circRPS19 knockdown (Fig. 2E and F). circRPS19 ablation led to a decrease in ECAR but an increase in OCR (Fig. 2G and H). ECAR/OCR ratio was decreased following circRPS19 interference (Fig. 2I). Taken together, these data indicated that circRPS19 facilitated GC cell proliferation and aerobic glycolysis.

*circRPS19 upregulates HK2 protein expression via USP7.*  A previous study identified three rate‑limiting enzymes in the regulation of aerobic glycolysis, including pyruvate kinases type M2 (PKM2), HK2 and phosphofructokinase 1 (PFK1) (14). Thus, the present study investigated which rate-limiting enzyme was regulated by circRPS19. RT-qPCR and western blotting showed that knockdown of circRPS19 had no effect on mRNA levels of these three enzymes (Fig. 3A) but effectively decreased protein levels of HK2 (Fig. 3B). Therefore, it was investigated whether circRPS19 influenced ubiquitination of HK2 protein. IP showed ubiquitination of



Figure 1. circRPS19 has circular characteristics and is upregulated in GC cells. (A) Differentially expressed circRNAs in GC tissue obtained from GSE83521 dataset. (B) Expression of five circRNAs previously unreported in GC were detected by RT-qPCR in AGS, HGC-27, NCI-N87, SNU-1 and GES-1 cells. (C) circRPS19 structure. (D) PCR‑agarose gel electrophoresis verified the circular structure of circRPS19 in GES‑1 cells. cDNA was created by RT of circRPS19; gDNA was the genome sequence of circRPS19. circRPS19 and RPS19 mRNA levels were detected by RT‑qPCR in GES‑1 cells following (E) RNase R and (F) ActD treatment. \*P<0.05, \*\*P<0.01 vs. control group. circRPS19, circRNA ribosomal protein S19; GC, gastric cancer; RT‑q, reverse transcription‑quantitative; n.s., not significant.

HK2 was notably increased by circRPS19 silencing (Fig. 3C). It was hypothesized circRPS19 may stabilize HK2 by directly interacting with it. However, RIP showed that circRPS19 did not directly bind to HK2 protein (Fig. 3D). The protein interacting with HK2 and modulating ubiquitination of HK2 protein was assessed by Co‑IP followed by mass spectrometry.



Figure 2. circRPS19 knockdown inhibits GC cell proliferation and aerobic glycolysis. (A) Proliferation of AGS and HGC‑27 cells following circRPS19 ablation was detected by Cell Counting Kit-8. (B) Proliferation of AGS and HGC-27 cells following circRPS19 ablation was detected by EdU assay. (C) TUNEL assay showed GC cell apoptosis following circRPS19 ablation (scale bar, 200 µm). (D) Flow cytometry assay showed GC cell apoptosis following circRPS19 ablation. (E) Glucose consumption, (F) lactate production, (G) ECAR and (H) OCR were determined in GC cells following circRPS19 silencing. (I) ECAR/OCR ratio was assessed in GC cells following circRPS19 silencing. \*P<0.05, \*\*P<0.01, \*\*\*P<0.001 vs. control group. circRPS19, circRNA ribosomal protein S19; GC, gastric cancer; ECAR, extracellular acidification rate; OCR, oxygen consumption rate; OD, optical density; si, small interfering; NC, negative control.



Figure 3. circRPS19 upregulates HK2 protein expression via USP7. (A) mRNA and (B) protein levels of PKM2, HK2 and PFK1 were assessed by RT‑qPCR and WB in AGS cells following circRPS19 ablation. (C) IP showed HK2 ubiquitination in AGS cells following circRPS19 depletion. (D) RIP assay was used to examine the interaction between circRPS19 and HK2 protein in AGS cells. (E) Co‑IP followed by mass spectrometry analysis was used in AGS cells to determine enzymes affecting ubiquitination of HK2 protein. (F) mRNA and (G) protein levels of USP7 were assessed by RT‑qPCR and WB in AGS cells following circRPS19 ablation. (H) FISH assay was used to determine co‑localization of circRPS19 and USP7 mRNA in AGS cells. (I) RIP assay was used to detect abundance of circRPS19 and USP7 mRNA in RISC. \*\*P<0.01, \*\*\*P<0.001, compared with control group. circRPS19, circRNA ribosomal protein S19; HK2, hexokinase 2; USP7, ubiquitin-specific processing protease 7; PKM2, pyruvate kinase M2; PFK1, 6-phosphofructokinase subunit alpha; RT-q, reverse transcription-quantitative; IP, immunoprecipitation; FISH, Fluorescence in situ hybridization; RISC, RNA-induced silencing complex; n.s., not significant; si, small interfering; NC, negative control; WB, western blot; IB, immunoblot; Ub, ubiquitin.

Deubiquitinating enzyme USP7 interacted with HK2 (Fig. 3E). RT‑qPCR and western blotting were performed in AGS cells to investigate if circRPS19 affected USP7 mRNA and protein expression. mRNA and protein levels of USP7 were significantly decreased by circRPS19 depletion (Fig. 3F and G). FISH assay was used to investigate co‑localization of circRPS19 and USP7 mRNA in AGS cells. circRPS19 and USP7 mRNA were co-localized in the cytoplasm (Fig. 3H). RIP showed that circRPS19 and USP7 mRNA were more abundant in AGO2 immunoprecipitate compared with that in IgG immunoprecipitate (Fig. 3I). In summary, circRPS19 upregulated USP7 expression to enhance HK2 protein levels and promote aerobic glycolysis of GC cells.

*CircRPS19 upregulates USP7 expression via competitively binding to miR-125a-5p.* It was hypothesized that circRPS19



Figure 4. circRPS19 upregulates USP7 expression via competitively binding to miR‑125a‑5p. (A) Common miRs interacting with USP7 and circRPS19 predicted by starBase database (starbase.sysu.edu.cn/). (B) RNA pulldown assay showed the combination of circRPS19 with miR-125a-5p, miR-125b-5p and miR-338-3p in AGS cells. (C) RNA pulldown assay revealed the interaction of miR-125a-5p with circRPS19 in AGS and HGC-27 cells. (D) RNA pulldown assay revealed the interaction of miR-125a-5p with USP7 3'-UTR in AGS and HGC-27 cells. (E) Luciferase reporter assay revealed the interaction of miR-125a-5p with circRPS19 in AGS and HGC-27 cells. (F) Luciferase reporter assay revealed the interaction of miR-125a-5p with USP7 3'UTR in AGS and HGC-27 cells. (G) miR-125a-5p expression was probed by RT-qPCR in AGS and HGC-27 cells subsequent to circRPS19 knockdown. (H) mRNA and (I) protein levels of USP7 were assessed by RT‑qPCR and western blot in AGS cells transfected with si‑NC, si‑circRPS19‑1, si‑circRPS19‑1 + inhibitor‑NC or si‑circRPS19‑1 + miR‑125a‑5p inhibitor. \*\*P<0.01, \*\*\*P<0.001, \*\*\*\*P<0.0001 compared with control group. circRPS19, circRNA ribosomal protein S19; USP7, ubiquitin-specific processing protease 7; miR, microRNA; UTR, untranslated region; RT-q, reverse transcription-quantitative; NC, negative control; bio, biotinylated; MUT, mutant; si, small interfering; n.s., not significant.



Figure 5. circRNA ribosomal protein S19 upregulates HK2 protein via USP7‑mediated deubiquitination. (A) Co‑IP analysis of the interaction between USP7 and HK2 in AGS cells. (B) Immunofluorescence assay detected the co-localization of USP7 and HK2 in AGS cells. WB of (C) HK2 protein expression in AGS cells and (D) degradation rate of HK2 protein in CHX‑treated AGS cells following USP7 ablation and (E) HK2 protein expression following USP7 overexpres‑ sion and HBX19818 treatment in AGS cells. (F) IP showed HK2 ubiquitination in MG132-treated AGS cells following USP7 depletion. \*P<0.05, \*\*P<0.01, \*\*P<0.001 vs. control group. HK2, hexokinase 2; USP7, ubiquitin-specific processing protease 7; IP, immunoprecipitation; CHX, cycloheximide; HBX19818, a specific inhibitor of USP7; si, small interfering; NC, negative control; WB, western blotting; Ub, ubiquitin.



Figure 6. circRPS19 facilitates GC cell proliferation and aerobic glycolysis by modulating USP7. (A) Cell Counting Kit-8 and (B) EdU assays showed proliferation in AGS and HGC-27 cells (scale bar=200  $\mu$ m). (C) TUNEL assay (scale bar=200  $\mu$ m) and (D) flow cytometry showed GC cell apoptosis. (E) Glucose consumption and (F) lactate production of GC cells. \*P<0.05, \*\*P<0.01 vs. control. circRPS19, circRNA ribosomal protein S19; GC, gastric cancer; OD, optical density; si, small interfering; NC, negative control.

might modulate USP7 via competitive endogenous RNA (ceRNA) network. starBase database was used to screen miRNAs that potentially interact with both circRPS19 and USP7. As a result, three common miRNAs (hsa-miR-125a-5p, hsa‑miR‑125b‑5p and hsa‑miR‑338‑3p) were obtained (Figs. S2A and B and 4A). RNA pulldown assay was used for screening in AGS cells and binding affinity of miR-125a-5p with circRPS19 was the highest compared with other miRNAs



Figure 7. circRPS19 promotes GC cell proliferation and aerobic glycolysis *in vivo*. (A) Tumor volume was recorded every 3 days. (B) Representative image and (C) weight of excised tumors. (D) Immunohistochemistry was implemented to investigate the expression of Ki-67, HK2 and USP7 in tumor tissues. (E) Glucose consumption and (F) lactate production were examined in tumor tissues. \*\*P<0.01 compared with control group. circRPS19, circRNA ribosomal protein S19; GC, gastric cancer; HK2, hexokinase 2; USP7, ubiquitin‑specific processing protease 7; si, small interfering; NC, negative control.

(Fig. 4B). RNA pulldown assay verified the interaction of miR-125a-5p with circRPS19 (Fig. 4C). RNA pulldown assay in AGS and HGC‑27 cells showed that miR‑125a‑5p could combine with USP7 3' UTR (Fig. 4D). Next, miR-125a-5p was overexpressed by transfection with miR‑125a‑5p mimics into AGS and HGC-27 cells and the overexpression efficiency were detected by RT‑qPCR (Fig. S1B). As indicated by luciferase reporter assay, upregulation of miR-125a-5p decreased luciferase activity of pmirGLO‑circRPS19/USP7‑3' UTR, while no obvious changes were observed in MUT groups. miR‑125a‑5p could bind with both circRPS19 and USP7 3' UTR (Fig. 4E and F). RT‑qPCR results indicated that circRPS19 silencing had no effects on miR-125a-5p expression (Fig. 4G), indicating that circRPS19 exerted its function by competitively binding to miR‑125a‑5p. mRNA and protein levels of USP7 suppressed by circRPS19 knockdown were completely reversed by miR‑125a‑5p inhibition (Fig. 4H and I). circRPS19 upregulated USP7 expression via competitively binding to miR-125a-5p.

*circRPS19 upregulates H2K protein through USP7‑mediated deubiquitination*. Based on the aforementioned interaction between USP7 and HK2, USP7‑mediated HK2 deubiquitination was investigated. Co-IP in AGS cells demonstrated an interaction between USP7 and HK2 (Fig. 5A). Immunofluorescence assay revealed that USP7 and HK2 were co-localized in the cytoplasm of AGS cells (Fig. 5B). To validate the regulatory effect of USP7 on HK2, USP7 expression was knocked down and efficiency was assessed by RT‑qPCR (Fig. S1C). HK2 protein was downregulated after USP7 interference (Fig. 5C). Degradation rate of HK2 protein was accelerated after interference with USP7 in CHX‑treated cells (Fig. 5D). The overexpression efficiency of pcDNA3.1‑USP7 was detected by RT‑qPCR in AGS and HGC‑27 cells (Fig. S1D). The western blot results showed that increased HK2 protein expression induced by USP7 overexpression was reversed by USP7 inhibitor HBX19818, indicating that USP7 mediated deubiquitination of HK2 (Fig. 5E). IP showed that HK2 ubiquitination was notably decreased by USP7 knockdown (Fig. 5F). circRPS19 promoted USP7‑mediated deubiquitination of HK2.

*circRPS19 facilitates GC cell proliferation and aerobic glycolysis by modulating USP7. In vitro* rescue assays were performed to investigate whether circRPS19 promotes aerobic glycolysis of GC cells by regulating USP7, thus facilitating GC cell progression. As indicated by CCK-8 and EdU assay, decreased GC cell proliferation induced by circRPS19 depletion was counteracted by USP7 overexpression (Fig. 6A and B). TUNEL and flow cytometry showed that the effect of circRPS19 knockdown on GC cell apoptosis was counteracted by USP7 overexpression (Fig. 6C and D). The present study investigated the roles of circRPS19 and USP7 in aerobic glycolysis of GC cells. Glucose consumption and lactate production suppressed by circRPS19 silencing was restored by USP7 overexpression (Fig. 6E and F). To summarize, circRPS19 facilitated proliferation and aerobic glycolysis but inhibited apoptosis of GC cells by modulating USP7.

*circRPS19 promotes GC cell proliferation and aerobic glycolysis in vivo.* The present study performed animal experi‑ ments to detect the effect of circRPS19 on GC *in vivo*. Tumor volume in nude mice was measured every 3 days beginning on day 7. The growth curve showed that circRPS19 knockdown decreased tumor growth (Fig. 7A). On day 28, tumors were resected. Size and volume of excised tumors were reduced in si-circRPS19-1 compared with si-NC group (Fig. 7B and C). Immunohistochemistry showed that the expression of Ki67, HK2, and USP7 was decreased in si-circRPS19-1 group compared with si‑NC group (Fig. 7D). The effects of circRPS19 on aerobic glycolysis of tumor cells were analyzed. Glucose consumption and lactate production of tumor cells were decreased in si-circRPS19-1 group in comparison with si-NC group (Fig. 7E and F). Therefore, circRPS19 promoted GC cell proliferation and aerobic glycolysis *in vivo*.

# **Discussion**

Previous studies have demonstrated that circRNAs play either a tumor‑suppressing or ‑promoting role in GC: For example, circRNA circ\_001988 can sequester miR‑197‑3p to hamper progression of GC both *in vitro* and *in vivo* (27); circRNA circ\_0004104 enhances GC progression by sequestering miR-539-3p and regulating ring finger protein 2 (RNF2) (28). In addition, circRNA Ran GTPase activating protein 1 (circ‑RanGAP1) targets miR‑877‑3p to modulate VEGFA expression, thereby promoting GC cell migration and invasion (29). The present study identified a novel circRNA, circRPS19, that was notably upregulated in GC cell lines. Furthermore, circRPS19 knockdown attenuated GC cell proliferation both *in vitro* and *in vivo*. FISH results demonstrated that circRPS19 and USP7 mRNA localized in the cytoplasm of GC cells. The cytoplasmic localization is indicative of the regulation of circRNA on mRNA at the post-transcriptional level (30). As indicated by previous studies, circRNAs serve as ceRNAs by competitively interacting with miRNAs to affect levels of target mRNAs: For example, circRNA DNA topoisomerase IIA as a ceRNA increases sushi domain containing 2 (SUSD2) expression via sequestering miR‑346 in glioma cells (31); circ\_0058357 competitively combines with miR‑24‑3p to increase AVL9 cell migration associated (AVL9) expression in non‑small cell lung cancer cells (32); circRNA itchy E3 ubiquitin protein ligase (circ‑ITCH) acts as a sponge for miR‑199a‑5p to enhance Klotho expression in GC cells (33). It was hypothesized that circRPS19 as a ceRNA binds to a certain miRNA to regulate USP7 mRNA. Through bioinformatics analysis, miR‑125a‑5p was screened. The regulatory influence of miR‑125a‑5p on GC progression has been studied previously (34). Additionally, tyrosine kinase non receptor 2 antisense RNA 1 (TNK2‑AS1) targets miR‑125a‑5p to modulate the PI3K/Akt pathway, thereby facilitating epithelial‑mesenchymal transition, invasion and migration of GC cells (35). The present study showed that miR-125a-5p bound to both circRPS19 and USP7 mRNA. Rescue assays proved that circRPS19 upregulated USP7 mRNA expression by competitively binding to miR-125a-5p.

Rate-limiting enzyme HK2 is a key factor in glycolysis of GC cells (36). According to previous studies, circRNA cullin 3 (circCUL3) upregulates transcription factor STAT3, which facilitates HK2 transcription to increase aerobic glycolysis of GC cells (19); circRNF20 increases aerobic glycolysis of breast cancer cells by upregulating hypoxia inducible factor-1 $\alpha$ , a transcription factor for HK2 (37). The present study showed that circRPS19 promoted aerobic glycolysis of GC cells by modulating HK2. Nevertheless, gene expression analysis showed that circRPS19 could only affect HK2 protein, excluding the possibility of ceRNA network and transcription activation. It was hypothesized that circRPS19 influences ubiquitination of HK2 protein via deubiquitinating enzyme USP7. USP7 mediates ubiquitination of target protein in cancer cells (38). The present study showed that circRPS19 can upregulate USP7 to mediate deubiquitination of HK2.

The present study showed that circRPS19 promotes GC cell proliferation via aerobic glycolysis. Furthermore, circRPS19 mediated deubiquitination of HK2 by USP7, which was also regulated by circRPS19 via ceRNA network. The present findings provide understanding of the molecular mechanisms underlying GC, which may facilitate development of treatment for GC.

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# **Availability of data and materials**

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

### **Authors' contributions**

SL and JQ conceived and designed the study. XZ and JS performed experiments. XZ analyzed and interpreted data. XZ and JQ wrote the manuscript. XZ and JS confirm the authenticity of all the raw data. All authors have read and approved the final manuscript.

## **Ethics approval and consent to participate**

The present study was reviewed and approved by the Ethics Committee of Affiliated Hospital of Nanjing University of Chinese Medicine (approval no. 2021DW‑35‑01; Nanjing, China).

## **Patient consent for publication**

Not applicable.

### **Competing interests**

The authors declare that they have no competing interests.

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